Medical Procedures
for MED-EL Implant Systems
This manual provides important instructions and safety information for MED-EL Implant System users who have to undergo a medical procedure (e.g. MRI).

As an implant user, you might have questions about undergoing further medical procedures. Your medical team may also want more information about any special considerations for implant users. This guidance provides information that will help prevent damage to your implant and injury to yourself. Please share this information with your healthcare provider.

Not all products in this document are currently approved or available in all countries. Please contact your local MED-EL representative for information on current product availability in your country.

In this document the general term “MED-EL Implant System” is used for all implant types. The specific implant name is identified in the header of the applicable section.
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Interference with other equipment, robustness of the device in special medical or diagnostic environments

- Instruments used in electrosurgery can produce high-frequency voltages which may induce currents in the electrodes of implantable devices. Such currents may damage the implant and/or the surrounding tissue. Monopolar electrosurgical instruments must not be used in the head and neck region. If bipolar electrosurgical instruments are used, the tips of the cautery must be kept at least 5 mm away from the reference electrodes on the stimulator housing and any contacts of the active electrode.
- Generally remove your external components (e.g. audio processor and accessories) from your head when undergoing medical treatment where an electrical current is passed through your body, or at least carefully observe the correct functioning of your entire MED-EL Implant System during the initial stages of the treatment.
- Any necessary ionising radiation therapy should be carefully considered and the risk of damage to the MED-EL implant has to be carefully weighed against the medical benefit of such therapy.
- Electroshock or electroconvulsive therapy in the head and neck region must not be used. Such therapy may damage the implant and/or the surrounding tissue.
- Neurostimulation or diathermy must not be carried out in the area of the implant since it could lead to current induction at the electrodes. This may damage the implant and/or the surrounding tissue. This applies also to iontophoresis and any current inducing medical and/or cosmetic treatment.
- Ultrasonic therapy and imaging must not be used in the area of the implant, as the implant may inadvertently concentrate the ultrasound field and cause harm.
- MED-EL implants are robust against radiotherapeutic irradiation of up to a total ionisation dose of 240 Gy. MED-EL external components need to be taken off during irradiation. Therapeutic ionising radiation in general may damage electronic components of your MED-EL Implant System and such damage may not be immediately detected. In order to minimise the risk of tissue necrosis due to local overdose, during radiotherapeutic treatments, the implant should not be placed in the direct radiotherapeutic beam.
- Other treatments: The effects of a number of treatments are unknown, e.g. electrical examinations in the dental area. Please contact your clinic.
MRI Caution

The external components of the MED-EL Implant System (audio processor and accessories) are MR Unsafe and need to be removed prior to scanning.

The implant components of the MED-EL Implant System are MR Conditional.

MRI is possible in patients with MED-EL implants only with specified models of MRI machines.

Evidence has been provided for this implant type to pose no known hazard in specified MRI environments (without surgical removal of the internal magnet) when adhering to the conditions and Safety Guidelines listed below. The implant has a specially designed magnet which allows safe MRI scanning with the magnet in place, and there is no need to remove the implant magnet regardless of the scanner field strength. The implant magnet can be surgically removed if needed to avoid imaging artefacts. The physician/MRI operator should always be informed that a patient is a MED-EL implant user and that special safety guidelines have to be followed.

MRI scanning is possible in consideration of the Safety Guidelines if the following conditions are fulfilled:

- MRI scanners with static magnetic fields of 0.2T, 1.0T, 1.5T or 3.0T only. No other field strengths are allowed. When using other field strengths, injury to the patient and/or damage to the implant are possible.
- In case of additional implants, e.g. a hearing implant in the other ear: MRI safety guidelines for this implant need to be considered in addition.
Safety Guidelines:
- Before patients enter any MRI room, all external components of the MED-EL Implant System (audio processor and accessories) must be removed from the head. An optional supportive head bandage may be placed over the implant. A supportive head bandage may be an elastic bandage wrapped tightly around the head at least three times (refer to Figure 1). The bandage shall fit tightly, but should not cause pain.
- For all MRI systems (1.0 T, 1.5 T and 3.0 T), the patient should be lying in the scanner in a supine, prone or side position with the head kept straight. The patient should be advised to not tilt their head to either side by more than 30 degrees from the long axis of the body otherwise torque will be exerted onto the implant magnet which might cause pain. In case of 0.2 T scanners, no specific head orientation is required.
- For 0.2T, 1.0T and 1.5T scans (see Table 1), only sequences in “Normal Operating Mode” with a maximum head Specific Absorption Rate (SAR) of 3.2 W/kg shall be used.
- For 3.0 T scans the SAR limit must not exceed the SAR values for specific anatomic regions given in Table 1 to avoid any potentially dangerous heating at the electrode contacts. For the same reason head transmit coils or multi-channel transmit coils must not be used in case of a 3.0 T MRI.

For head scans and scans with a landmark location that is less than 35 cm from the top of the head the MRI system must be able to provide a SAR limit prediction that allows fractional SAR display.

Sequences in Normal Operating Mode only with the following SAR restrictions:
- For head scans: Maximum average head SAR must not exceed 1.6 W/kg (50 % of maximum head SAR).
- For landmark locations less than 35 cm from the top of the head: Maximum whole-body SAR must not exceed 1.0 W/kg.
- For landmark locations at least 35 cm away from the top of the head: Maximum whole-body SAR must not exceed 2.0 W/kg.

<table>
<thead>
<tr>
<th>MRI field strengths</th>
<th>Average head SAR</th>
<th>Average whole-body SAR</th>
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<tr>
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<td>Landmark location &lt;35 cm from the top of the head</td>
<td>Landmark location ≥35 cm from the top of the head</td>
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<td>0.2T</td>
<td>3.2 W/kg</td>
<td>2.0 W/kg</td>
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<td>1.0T</td>
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<td>1.5T</td>
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<td>3.0T</td>
<td>1.6 W/kg</td>
<td>1.0 W/kg</td>
</tr>
</tbody>
</table>

Table 1: Specific Absorption Rate (SAR levels)
• During the scan patients might perceive auditory sensations such as clicking or beeping. Adequate counselling of the patient is advised prior to performing the MRI. The likelihood and intensity of auditory sensations can be reduced by selecting sequences with a lower Specific Absorption Rate (SAR) and slower gradient slew rates.

• The magnet can be surgically removed by pushing on the top side of the magnet so that it comes out at the bottom side of the implant to reduce image artefacts. If the magnet is not removed, image artefacts are to be expected (refer to Figure 2 and Figure 3).

• The exchange of the magnets with the Non-Magnetic Spacer and vice versa has been tested for at least five repetitions.

• The above instructions should also be followed if areas of the body other than the head are to be examined (e.g. knee, etc.). When lower extremities are to be examined, it is recommended that the patient’s legs are positioned in the scanner first.

If the conditions for MRI safety and the Safety Guidelines are not followed, injury to the patient and/or damage to the implant may result!
Figure 1: Head bandage to support fixation of the implant

Figure 2: Image artefacts arising in a 1.5T scanner. The left picture shows the artefacts obtained with the implant magnet in place, whereas the right picture illustrates the image artefacts when the implant magnet is replaced with the Non-Magnetic Spacer.

Figure 3: Image artefacts arising in a 3.0T scanner. The left picture shows the artefacts obtained with the implant magnet in place, whereas the right picture illustrates the image artefacts when the implant magnet is replaced with the Non-Magnetic Spacer.
Interference with other equipment, robustness of the device in special medical or diagnostic environments

- Instruments used in electrosurgery can produce high-frequency voltages which may induce currents in the electrodes of implantable devices. Such currents may damage the implant and/or the surrounding tissue. Monopolar electrosurgical instruments must not be used in the head and neck region. If bipolar electrosurgical instruments are used, the tips of the cautery must be kept at least 5 mm away from the reference electrodes on the stimulator housing and any contacts of the active electrode.
- Generally remove your external components (e.g. audio processor and accessories) from your head when undergoing medical treatment where an electrical current is passed through your body, or at least carefully observe the correct functioning of your entire MED-EL Implant System during the initial stages of the treatment.
- Any necessary ionising radiation therapy should be carefully considered and the risk of damage to the MED-EL implant has to be carefully weighed against the medical benefit of such therapy.
- Electroshock or electroconvulsive therapy in the head and neck region must not be used. Such therapy may damage the implant and/or the surrounding tissue.
- Neurostimulation or diathermy must not be carried out in the area of the implant since it could lead to current induction at the electrodes. This may damage the implant and/or the surrounding tissue. This applies also to iontophoresis and any current inducing medical and/or cosmetic treatment.
- Ultrasonic therapy and imaging must not be used in the area of the implant, as the implant may inadvertently concentrate the ultrasound field and cause harm.
- MED-EL implants are robust against radiotherapeutic irradiation of up to a total ionisation dose of 240 Gy. MED-EL external components need to be taken off during irradiation. Therapeutic ionising radiation in general may damage electronic components of your MED-EL Implant System and such damage may not be immediately detected. In order to minimise the risk of tissue necrosis due to local overdose, during radiotherapeutic treatments, the implant should not be placed in the direct radiotherapeutic beam.
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- MRI scanners with static magnetic fields of 0.2T, 1.0T or 1.5T only. No other field strengths are allowed. When using other field strengths, injury to the patient and/or damage to the implant are possible.
- In case of additional implants, e.g. a hearing implant in the other ear: MRI safety guidelines for this implant need to be considered in addition.
Safety Guidelines:

- Before patients enter any MRI room, all external components of the MED-EL Implant System (audio processor and accessories) must be removed from the head. An optional supportive head bandage may be placed over the implant. A supportive head bandage may be an elastic bandage wrapped tightly around the head at least three times (refer to Figure 1). The bandage shall fit tightly, but should not cause pain.

- For all MRI systems (1.0 T, 1.5 T), the patient should be lying in the scanner in a supine, prone or side position with the head kept straight. The patient should be advised to not tilt their head to either side by more than 30 degrees from the long axis of the body otherwise torque will be exerted onto the implant magnet which might cause pain. In case of 0.2 T scanners, no specific head orientation is required.

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- During the scan patients might perceive auditory sensations such as clicking or beeping. Adequate counselling of the patient is advised prior to performing the MRI. The likelihood and intensity of auditory sensations can be reduced by selecting sequences with a lower Specific Absorption Rate (SAR) and slower gradient slew rates.

- The magnet can be surgically removed by pushing on the top side of the magnet so that it comes out at the bottom side of the implant to reduce image artefacts. If the magnet is not removed, image artefacts are to be expected (refer to Figure 2).

- The exchange of the magnets with the Non-Magnetic Spacer and vice versa has been tested for at least five repetitions.

- The above instructions should also be followed if areas of the body other than the head are to be examined (e.g. knee, etc.). When lower extremities are to be examined, it is recommended that the patient's legs are positioned in the scanner first.

If the conditions for MRI safety and the Safety Guidelines are not followed, injury to the patient and/or damage to the implant may result!
Figure 1: Head bandage to support fixation of the implant

Figure 2: Image artefacts arising in a 1.5T scanner. The left picture shows the artefacts obtained with the implant magnet in place, whereas the right picture illustrates the image artefacts when the implant magnet is replaced with the Non-Magnetic Spacer.
Interference with other equipment, robustness of the device in special medical or diagnostic environments

- Instruments used in electrosurgery can produce high-frequency voltages which may induce currents in the electrodes of implantable devices. Such currents may damage the implant and/or the surrounding tissue. Monopolar electrosurgical instruments must not be used in the head and neck region. If bipolar electrosurgical instruments are used, the tips of the cautery must be kept at least 5 mm away from the reference electrodes on the stimulator housing and any contacts of the active electrode.
- Generally remove your external components (e.g. audio processor and accessories) from your head when undergoing medical treatment where an electrical current is passed through your body, or at least carefully observe the correct functioning of your entire MED-EL Implant System during the initial stages of the treatment.
- Any necessary ionising radiation therapy should be carefully considered and the risk of damage to the MED-EL implant has to be carefully weighed against the medical benefit of such therapy.
- Electroshock or electroconvulsive therapy in the head and neck region must not be used. Such therapy may damage the implant and/or the surrounding tissue.
- Neurostimulation or diathermy must not be carried out in the area of the implant since it could lead to current induction at the electrodes. This may damage the implant and/or the surrounding tissue. This applies also to iontophoresis and any current inducing medical and/or cosmetic treatment.
- Ultrasonic therapy and imaging must not be used in the area of the implant, as the implant may inadvertently concentrate the ultrasound field and cause harm.
- MED-EL implants are robust against radiotherapeutic irradiation of up to a total ionisation dose of 240 Gy. MED-EL external components need to be taken off during irradiation. Therapeutic ionising radiation in general may damage electronic components of your MED-EL Implant System and such damage may not be immediately detected. In order to minimise the risk of tissue necrosis due to local overdose, during radiotherapeutic treatments, the implant should not be placed in the direct radiotherapeutic beam.
- Other treatments: The effects of a number of treatments are unknown, e.g. electrical examinations in the dental area. Please contact your clinic.
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MRI is possible in patients with MED-EL implants only with specified models of MRI machines.

Evidence has been provided for these implants to pose no known hazard in magnetic field strengths of 0.2T, 1.0T and 1.5T (without surgical removal of the internal magnet) when adhering to the following safety recommendations and guidelines. The physician/MRI operator should always be informed that a patient is a MED-EL implant user and that special safety recommendations and guidelines have to be followed.

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• In case of additional implants, e.g. a hearing implant in the other ear: MRI safety guidelines for this implant need to be considered in addition.
Safety Guidelines:

- Before patients enter any MRI room, all external components of the MED-EL Implant System (audio processor and accessories) must be removed. For field strengths of 1.0 T or 1.5 T, a supportive head bandage must be placed over the implant. A supportive head bandage may be an elastic bandage wrapped tightly around the head at least three times (refer to Figure 1). The bandage shall fit tightly, but should not cause pain.
- In 1.0 T and 1.5 T MRI systems, the patient should be lying in the scanner in a supine, prone or side position with the head kept straight. The patient should be advised to not tilt their head to either side otherwise demagnetisation of the implant magnet may be possible. In case of 0.2 T scanners, no specific head orientation is required.
- Only sequences in Normal Operating Mode shall be used!
- During the scan patients might perceive auditory sensations such as clicking or beeping. Adequate counselling of the patient is advised prior to performing the MRI. The likelihood and intensity of auditory sensations can be reduced by selecting sequences with a lower Specific Absorption Rate (SAR) and slower gradient slew rates.
- Image artefacts are to be expected (refer to Figure 2).
- The above instructions should also be followed if areas of the body other than the head are to be examined (e.g. knee, etc.). When lower extremities are to be examined, it is recommended that the patient’s legs are positioned in the scanner first to minimise any risk of weakening the implant magnet.

If the conditions for MRI safety and the Safety Guidelines are not followed, injury to the patient and/or damage to the implant may result!
Figure 1: Head bandage to support fixation of the implant

Figure 2: MR images obtained with a 1.5T scanner (8-year-old child)
Interference with other equipment, robustness of the device in special medical or diagnostic environments

• Instruments used in electrosurgery can produce high-frequency voltages which may induce currents in the electrodes of implantable devices. Such currents may damage the implant and/or the surrounding tissue. Monopolar electrosurgical instruments must not be used in the head and neck region. If bipolar electrosurgical instruments are used, the tips of the cautery must be kept at least 5 mm away from the reference electrodes on the stimulator housing and any contacts of the active electrode.
• Generally remove your external components (e.g. audio processor and accessories) from your head when undergoing medical treatment where an electrical current is passed through your body, or at least carefully observe the correct functioning of your entire MED-EL Implant System during the initial stages of the treatment.
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• Ultrasonic therapy and imaging must not be used in the area of the implant, as the implant may inadvertently concentrate the ultrasound field and cause harm.
• MED-EL implants are robust against radiotherapeutic irradiation of up to a total ionisation dose of 240 Gy. MED-EL external components need to be taken off during irradiation. Therapeutic ionising radiation in general may damage electronic components of your MED-EL Implant System and such damage may not be immediately detected. In order to minimise the risk of tissue necrosis due to local overdose, during radiotherapeutic treatments, the implant should not be placed in the direct radiotherapeutic beam.
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- In case of additional implants, e.g. a hearing implant in the other ear: MRI safety guidelines for this implant need to be considered in addition.
Safety Guidelines:

- Before patients enter any MRI room, all external components of the MED-EL Implant System (audio processor and accessories) must be removed. For field strengths of 1.0 T or 1.5 T, a supportive head bandage must be placed over the implant. A supportive head bandage may be an elastic bandage wrapped tightly around the head at least three times (refer to Figure 1). The bandage shall fit tightly, but should not cause pain.
- In 1.0 T and 1.5 T MRI systems, the patient should be lying in the scanner in a supine, prone or side position with the head kept straight. The patient should be advised to not tilt their head to either side otherwise demagnetisation of the implant magnet may be possible. In case of 0.2 T scanners, no specific head orientation is required.
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- Image artefacts are to be expected (refer to Figure 2).
- The above instructions should also be followed if areas of the body other than the head are to be examined (e.g. knee, etc.). When lower extremities are to be examined, it is recommended that the patient’s legs are positioned in the scanner first to minimise any risk of weakening the implant magnet.

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- Generally remove your external components (e.g. audio processor and accessories) from your head when undergoing medical treatment where an electrical current is passed through your body, or at least carefully observe the correct functioning of your entire MED-EL Implant System during the initial stages of the treatment.
- Any necessary ionising radiation therapy should be carefully considered and the risk of damage to the MED-EL implant has to be carefully weighed against the medical benefit of such therapy.
- Electroshock or electroconvulsive therapy in the head and neck region must not be used. Such therapy may damage the implant and/or the surrounding tissue.
- Neurostimulation or diathermy must not be carried out in the area of the implant since it could lead to current induction at the electrodes. This may damage the implant and/or the surrounding tissue. This applies also to iontophoresis and any current inducing medical and/or cosmetic treatment.
- A diagnostic level of ultrasonic energy of up to 500 W/m² within the range of 2 MHz to 5 MHz does not cause any damage to the implant.
- MED-EL implants are robust against radiotherapeutic irradiation of up to a total ionisation dose of 240 Gy. MED-EL external components need to be taken off during irradiation. Therapeutic ionising radiation in general may damage electronic components of your MED-EL Implant System and such damage may not be immediately detected. In order to minimise the risk of tissue necrosis due to local overdose, during radiotherapeutic treatments, the implant should not be placed in the direct radiotherapeutic beam.
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• MRI scanners with static magnetic field strengths of 0.2T, 1.0T or 1.5T only. No other field strengths are allowed. When using other field strengths, injury to the patient and/or damage to the implant are possible.
• MRI scan not earlier than 6 months post implantation. Performing a MRI at an earlier stage may result in implant displacement and/or damage to the implant.
• A minimum thickness of the bone underneath the implant magnet of 0.4 mm is required to withstand forces of 5 N (equals a gravitational force of about 0.5 kg). In a MRI scanner torque forces act on the implant magnet, exerting rotational pressure: the device will try to turn to line up with force lines. The resulting forces on the edges of the implant are counterbalanced by the cranial bone and the skin flap. The bone underneath the implant magnet should be thick enough to withstand these exerting forces.
• Patients with mechanically damaged implants must not undergo MRI. Ignoring this guideline could result in injury to the patient.
Safety Guidelines:

- Before patients enter any MRI room, all external components of the MED-EL Implant System (audio processor and accessories) must be removed. For field strengths of 1.0 T or 1.5 T, a supportive head bandage must be placed over the implant. A supportive head bandage may be an elastic bandage wrapped tightly around the head at least three times (refer to Figure 1). The bandage shall fit tightly, but should not cause pain.
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- The above instructions also apply to patients with bilateral MED-EL implants.

If the conditions for MRI safety and the Safety Guidelines are not followed, injury to the patient and/or damage to the implant may result!
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Figure 2: MR images obtained with a 1.5T scanner (8-year-old child)
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- Generally remove your external components (e.g. audio processor and accessories) from your head when undergoing medical treatment where an electrical current is passed through your body, or at least carefully observe the correct functioning of your entire MED-EL Implant System during the initial stages of the treatment.

- Any necessary ionising radiation therapy should be carefully considered and the risk of damage to the MED-EL implant has to be carefully weighed against the medical benefit of such therapy.

- Electroshock or electroconvulsive therapy in the head and neck region must not be used. Such therapy may damage the implant and/or the surrounding tissue.

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- MRI scanners with static magnetic field strengths of 0.2T, 1.0T or 1.5T only. No other field strengths are allowed. When using other field strengths, injury to the patient and/or damage to the implant are possible.
- MRI scan not earlier than 6 months post implantation. Performing a MRI at an earlier stage may result in implant displacement and/or damage to the implant.
- A minimum thickness of the bone underneath the implant magnet of 0.4mm is required to withstand forces of 5 N (equals a gravitational force of about 0.5 kg) or up to 9 N for the C40 cochlear implant. In a MRI scanner torque forces act on the implant magnet, exerting rotational pressure: the device will try to turn to line up with force lines. The resulting forces on the edges of the implant are counterbalanced by the cranial bone and the skin flap. The bone underneath the implant magnet should be thick enough to withstand these exerting forces.
- Patients with mechanically damaged implants must not undergo MRI. Ignoring this guideline could result in injury to the patient.
Safety Guidelines:

• Before patients enter any MRI room, all external components of the MED-EL Implant System (audio processor and accessories) must be removed. For field strengths of 1.0 T or 1.5 T, a supportive head bandage must be placed over the implant. A supportive head bandage may be an elastic bandage wrapped tightly around the head at least three times (refer to Figure 1). The bandage shall fit tightly, but should not cause pain.

• In 1.0 T and 1.5 T MRI systems, the patient should be lying in the scanner in a supine, prone or side position with the head kept straight. The patient should be advised to not tilt their head to either side otherwise demagnetisation of the implant magnet may be possible. In case of 0.2 T scanners, no specific head orientation is required.

• Only sequences in Normal Operating Mode shall be used!

• During the scan patients might perceive auditory sensations such as clicking or beeping. Adequate counselling of the patient is advised prior to performing the MRI. The likelihood and intensity of auditory sensations can be reduced by selecting sequences with a lower Specific Absorption Rate (SAR) and slower gradient slew rates.

• Image artefacts are to be expected (refer to Figure 2).

• The above instructions should also be followed if areas of the body other than the head are to be examined (e.g. knee, etc.). When lower extremities are to be examined, it is recommended that the patient's legs are positioned in the scanner first to minimise any risk of weakening the implant magnet.

• The above instructions also apply to patients with bilateral MED-EL implants.

If the conditions for MRI safety and the Safety Guidelines are not followed, injury to the patient and/or damage to the implant may result!
Figure 1: Head bandage to support fixation of the implant

Figure 2: MR images obtained with a 1.5T scanner (8-year-old child)
Help and assistance are always available from your local office. Please refer to the accompanying Contact Sheet for your local office.

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